OXALICUM ACIDUM- oxalic acid dihydrate liquid Washington Homeopathic Products

Disclaimer: This homeopathic product has not been evaluated by the Food and Drug Administration for safety or efficacy. FDA is not aware of scientific evidence to support homeopathy as effective.

DRUG FACTS

ACTIVE INGREDIENTS

OXALICUM AC

USES

To relieve the symptoms of weak digestion.

KEEP OUT OF REACH OF CHILDREN

Keep this and all medicines out of reach of children.

INDICATIONS

Indications:

OXALICUM AC Weak digestion

STOP USE AND ASK DOCTOR

If symptoms persist/worsen or if pregnant/nursing, consult your practitioner.

DIRECTIONS

Adults: 4 drops into a tsp. of water 3 times a day. Children: 1/2 dose. Repeat at greater intervals as condition subsides. Or as directed by a lic. practitioner.

INACTIVE INGREDIENTS

Sucrose/Lactose

PRINCIPAL DISPLAY PANEL

The OTC potency range of OXALICUM AC is 6x–30x, 3c–30c, 200c, 1m, 10m, 50m, and CM.

Availability is subject to change.



All WHP single remedies are made to order; thus, the labels are printed on the same label stock, as the orders are filled.

'Bottle Size,' 'Potency,' and 'Alcohol Percentage' vary on the label depending on customer choice. Standard bottle sizes for dilution-form remedies are 15ml, 30ml, 50ml, and 100ml.

OXALICUM ACIDUM

oxalic acid dihydrate liquid

Product Information

Product Type	HUMAN OTC DRUG	Item Code (Source)	NDC:71919-513

Route of Administration ORAL

Active Ingredient/Active Moiety

Ingredient Name	Basis of Strength	Strength
OXALIC ACID DIHYDRATE (UNII: 0K2L2IJ59O) (OXALIC ACID -	OXALIC ACID DIHYDRATE	30 [hp_C] in 1 mL

Inactive Ingredients

Ingredient Name	Strength
ALCOHOL (UNII: 3K9958V90M)	
WATER (UNII: 059QF0KO0R)	

Product Characteristics

Color	white (white)	Score	
Shape		Size	
Flavor		Imprint Code	
Contains			

Packaging

#	Item Code	Package Description	Date	Marketing End Date
1	NDC:71919-513- 07	15 mL in 1 VIAL, GLASS; Type 0: Not a Combination Product	03/11/2010	
2	NDC:71919-513- 08	30 mL in 1 VIAL, GLASS; Type 0: Not a Combination Product	03/11/2010	
3	NDC:71919-513- 09	50 mL in 1 BOTTLE, GLASS; Type 0: Not a Combination Product	0 3/11/20 10	
4	NDC:71919-513- 10	100 mL in 1 BOTTLE, GLASS; Type 0: Not a Combination Product	03/11/2010	

Marketing Information			
Marketing Category	Application Number or Monograph Citation	Marketing Start Date	Marketing End Date
unapproved homeopathic		03/11/2010	

Labeler - Washington Homeopathic Products (084929389)

Establishment				
Name	Address	ID/FEI	Business Operations	
Washington Homeopathic Products		084929389	manufacture(71919-513)	

Revised: 12/2018 Washington Homeopathic Products